RESPONSE AND REQUEST FOR RECONSIDERATION

In response to the Office Action of July 18, 2006, Applicants hereby request the Examiner to reconsider the claims in view of the present amendments and remarks.

Applicants have amended independent claims 1 and 19 by specifying that the phosphite is defined as an alkyl phosphite rather than a hydrocarbyl phosphite as previously claimed. Support for this amendment is found in the specification on paragraph 18. This amendment serves to focus the claims on those specific phosphites that contain alkyl groups, rather than other, more broadly defined, hydrocarbyl groups.

Dependent claims 2 and 17 have been amended to specify that the phosphite definition is consistent with independent claims 1 and 19. Specifically, claims 2 and 17 specify the phosphite is an alkyl phosphite.

Applicants have added a new independent claim 23 and new dependent claims 24 and 25. Each of these claims is substantially the same as independent claim 1, except the alkyl phosphite is defined as "an alkyl phosphite, wherein the alkyl group contains 14 to 20 carbon atoms" or "12 to 30 carbon atoms." Support for the phosphite definition specifying the alkyl group contains 14 to 20 or 12 to 30 carbon atoms is found in paragraph 18 of the specification. No elements other than the nature and chain length of the alkyl groups have been amended.

Applicants submit that the amended claims are fully supported by the specification and do not add subject matter. Moreover, the claims as amended are novel and non-obvious over the cited prior art.

Applicants note with gratitude that the Examiner has withdrawn all objections and rejections, except those described in the office action dated July 18, 2006.

The Examiner, however, maintained that claims 1, 2, 4, 5, 8-10, 14 and 16-22 still failed to meet the requirements of 35 U.S.C. 103(a) over Sumiejski and Vinci et al. Further the Examiner maintained that claims 6, 7 and 15 still failed to meet the requirements of 35 U.S.C. 103(a) over Sumiejski and Vinci et al. and Tagliamonte.

The Examiner was of the position that the evidence supplied in the declaration by J. Sumiejski on May 1, 2006, was insufficient to establish unexpected or surprising results for the claimed invention. In particular, the data was deemed insufficient to rebut a prima facie case of obviousness since the data was believed to be not commensurate in scope with the claimed invention. Applicants respectfully traverse the Examiner's position.

Applicants submit that the experimental data provided in the declaration by Sumiejski is commensurate in scope with the invention as it is presently defined by the claims. In particular, Applicants presented data from an alkyl phosphite with a C14 alkyl group, as it provides a meaningful comparison of the lower limit of the present invention, presently set at C12, with the prior art. A person skilled in the art would expect that a C12-alkyl phosphite and the C14-alkyl phosphite actually tested would have similar performance because both alkyl chains have similar properties. As an example of this similarity, the

alcohols which correspond to these and longer alkyl chains are solids at around ambient temperature (about 23 °C). In contrast, alcohols with alkyl chains with 10 or fewer carbon atoms are typically liquids. The attached pages from the CRC Handbook of Chemistry and Physics, 75th Edition illustrates the melting points for decanol, dodecanol and tetradecanol. It will also be recognized that the longer chain alkyl groups share the property of increasing oil solubility, which is important for a composition designed for use as a lubricant.

It is therefore believed that properties of alkyl chains of 12 and more carbon atoms are sufficiently different from those in the reference, below 12, that the material having a C14 alkyl chain reasonably represents the class of materials with longer chain alkyl groups. As a consequence, while the declaration compared a C14-alkyl phosphite with the C6-alkyl phosphite of the reference, similar performance advantages would be expected from alkyl groups such as C12, or C14, or C16, C18, C20 or higher alkyl groups. Hence the invention example demonstrated in the declaration is commensurate with the scope of the alkyl phosphite of the invention as defined in claim 1, in terms of the nature and length of the alkyl groups.

The C14 test sample even more clearly supports the patentability of the narrower ranges of carbon atoms set forth in new claims 23, 24, and 25, and separate consideration for the subject matter of these claims is respectfully requested.

Accordingly, Applicants request the Examiner to find the experimental data provided to be commensurate with the scope of the independent claims and find all claims allowable.

With regard to the example presented in the declaration, the Examiner has further indicated that only one type of each component (b), (c) and (d) is exemplified. Applicants respectfully submit that further experimental exemplification of these components is not legally required. Components (b), (c), and (d), while required components of the lubricating composition, are only of secondary importance when considering the technical problem of reducing wear and shudder that is solved principally by component (a), i.e., the alkyl phosphite. And in order to determine whether an invention exhibits unexpected advantages over a prior art reference, it is only required to compare the claimed invention with the closest teaching of the prior art. The proper comparison of the prior art with a lubricating composition of the present invention is one that requires the fewest changes to the lubricating composition. This minimizes the number of variables altered and most clearly demonstrates the advantages of the invention. Applicants have closely reviewed the prior art and selected the closest explicit teaching (i.e., an example) to the present invention.

It is not legally necessary to compare variants of the claimed invention with hypothetical variations of the prior art that are not in fact disclosed. Thus, it is not required to test samples with variations of (b), (c), and (d) against modifications of the prior art in which (b), (c), and (d) are similarly varied. If Applicants were required to compare the present invention against such hypothetical, imagined, or reconstructed prior art compositions, it would not be a valid or useful or legally meaningful comparison. This is in part because

multiple variables would be changed, and in part because the comparison would no longer be against the actual prior art. Therefore, Applicants' obligation is to compare the invention only against the closest prior art and not against hypothetical, imagined, or reconstructed prior art. Accordingly, Applicants respectfully request the Examiner to withdraw objections of improper breadth of components (b), (c) and (d).

For the foregoing reasons it is submitted that the present claims are unobvious and in condition for allowance. The foregoing remarks are believed to be a full and complete response to the outstanding office action. Therefore an early and favorable reconsideration is respectfully requested. If the Examiner believes that only minor issues remain to be resolved, a telephone call to the Undersigned is suggested.

The number of claims, after amendment, is increased from the original 22 to 24. Please charge the fee for the additional 2 claims, believed to be $2 \times \$50 = \100.00 , to deposit account 12-2275 (The Lubrizol Corporation). The number of independent claims is now 3, for which no additional fee is due. Any additional required fees, or any insufficiency or overpayment of fees, should be charged or credited to this same account.

Respectfully submitted,

/ David M. Shold # 31664 /

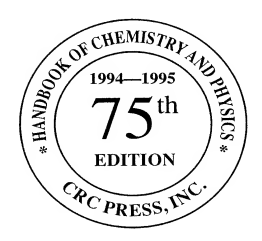
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CRC Press

Boca Raton Ann Arbor London Tokyo

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No claim to original U.S. Government works
International Standard Book Number 0-8493-0475-X
Library of Congress Card Number 13-11056
Printed in the United States of America 1 2 3 4 5 6 7 8 9 0
Printed on acid-free paper

PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS (continued)

No.	Name Synonym	Mol. Form. Mol. Wt.	CAS RN mp/°C	Merck No. bp/°C	Beil. Ref. den/g cm ⁻³	Solubility n _D
5066	Decanoic acid, methyl ester	C ₁₁ H ₂₂ O ₂ 186.29	110-42-9 -18	224	4-02-00-01044 0.8730 ²⁰	H ₂ O 1; EtOH 4; eth 4; ctc
5067	Decanoic acid, 1-methylethyl ester	C ₁₃ H ₂₈ O ₂ 214.35	2311-59-3	12110	4-02-00-01045 0.8543 ²⁰	
5068	Decanoic acid, 2-octyl-	C ₁₈ H ₃₈ O ₂	619-39-6		4-02-00-01254	1.4221 ²⁵ eth 4; EtOH 4
5069	Decanoic acid, propyl ester	284.48 C ₁₃ H ₂₆ O ₂	38.5 30673-60-0	215 ¹³	0.8447 ⁷⁰ 4-02-00-01045	
5070	1-Decanol	214.35 ² C ₁₀ H ₂₂ O	112-30-1	128.5 ¹⁰ 2847	0.8623 ²⁰ 4-01-00-01815	1.4280 ²⁰
	Capric alcohol 2-Decanol, (±)-	158.28	6.9	231.1	0.829720	H ₂ O 1; EtOH 5; eth 5; ace 1.4372 ²⁰
071	2-Decanol (DL)	C ₁₀ H ₂₂ O 158.28	74742-10-2 -2.4	211	4-01-00-01823 0.8250 ²⁰	EtOH 3; eth 5; ace 5; bz 3 1,4326 ²⁵
072	4-Decanol 1-Propylheptyl alcohol	C ₁₀ H ₂₂ O 158.28	2051-31-2 -11	210.5	4-01-00-01824 0.8261 ²⁰	H ₂ O 1; EtOH 3; ctc 3 1.4320 ²⁰
073	1-Decanol, 10-chloro-	C ₁₀ H ₂₁ CIO 192.73	51309-10-5	185-9 ¹⁵	4-01-00-01821	eth 4; EtOH 4
074	1-Decanol, 10-fluoro-	C10H21FO	12.5 334-64-5		0.9630 ²⁵ 4-01-00-01821	1.4578 ²⁰ eth 4; EtOH 4
075	10-Fluoro-1-decanol 2-Decanone	176.27 C10H20O	22 693-54-9	136-7 ¹⁵	0.919 ²⁰ 4-01-00-03367	1.4322 ²⁵ . H ₂ O 1; EtOH 3; eth 3; ctc
	Methyl octyl ketone 3-Decanone	C ₁₀ H ₂₀ O 156.27	14 928-80-3	210; 96 ¹²	0.8248 ²⁰	1.4255 ²⁰
076	Ethyl heptyl ketone	C ₁₀ H ₂₀ O 156.27	2.5	203	4-01-00-03368 0.8251 ²⁰	EtOH 3; eth 3; ctc 3 1.4252 ²⁰
077	4-Decanone Hexyl propyl ketone	C ₁₀ H ₂₀ O 156.27	624-16-8 -9	206.5	4-01-00-03368 0.824 ²⁰	H ₂ O 1; EtOH 5; eth 5 1.4240 ²¹
078	Decanoyl chloride Caprinoyl chloride	C ₁₀ H ₁₉ CIO 190.71	112-13-0		4-02-00-01050	eth 3; ctc 3
079	Decasiloxane, docosamethyl-	C ₂₂ H ₈₈ O ₉ Si ₁₀ 755.62	-34.5 556-70-7	95	0.919 ²⁵ 3-04-00-01881	1.4410 ²⁰ bz 4; lig 4
080	Decasiloxane, dicosamethyl 2-Decenal	755.62 CapHaeO	3913-71-1	183.4	0.925 ²⁰ 4-01-00-03511	1.398820
081	3-Decenal	C ₁₀ H ₁₈ O 154.25		230	0.845 ¹⁷	1.4533 ¹⁷
		C ₁₀ H ₁₈ O 154.25	58474-80-9	93-414	4-01-00-03512 0.850 ¹⁵	1.4462 ¹⁵
082	1-Decene	C ₁₀ H ₂₀ 140.27	872-05-9 -66.3	170.5	3-01-00-00858 0.7408 ²⁰	H ₂ O 1; EtOH 5; eth 5 1,4215 ²⁰
383	4-Decene	C ₁₀ H ₂₀ 140.27	19689-18-0		4-01-00-00902	
084	5-Decene, (É)-	C ₁₀ H ₂₀ 140.27	7433-56-9	170.6	0.7404 ²⁰ 4-01-00-00902	1.4243 ²⁰ H ₂ O 1; EtOH 5; eth 5; ctc
085	5-Decene, (Z)-	140.27 C ₁₀ H ₂₀	-73 7433-78-5	171	0.7401 ²⁰ 3-01-00-00859	1.4243 ²⁰ H ₂ O 1, EtOH 5, eth 5, ctc
086	1-Decene, 2-bromo-	140.27	-112	171; 73 ²⁰	0.7445 ²⁰	1.425820
-	2-Bromo-1-decene	C ₁₀ H ₁₉ Br 219.16	3017-67-2	115-6 ²²	3-01-00-00859 1.0844 ²⁰	1.4629 ²⁰
087	2-Decene, 1-bromo- 1-Bromo-2-decene	C ₁₀ H ₁₉ Br 219.16	14304-30-4	121 ¹⁷	4-01-00-00902 1,074 ¹⁸	lig 4 1.4716 ¹⁸
880	2-Decenoic acid Δ 2-Decenoic acid	C ₁₀ H ₁₈ O ₂	3913-85-7		4-02-00-01606	
089	3-Decenoic acid	170.25 C ₁₀ H ₁₈ O ₂	12 15469-77-9	165 ¹⁵	0.9280 ¹⁸ 4-02-00-01606	1.4616 ²⁰
090	4-Decenoic acid	170.25 C ₁₀ H ₁₈ O ₂	18 26303-90-2	154-63 ¹¹	0.914 ¹⁵ 4-02-00-01607	1.4510 ¹⁸ bz 4; eth 4
091	Deconic acid telomer	170.25		14913	0.919720	1.4497 ²⁰
	9-Decenoic acid Caproleic acid	C ₁₀ H ₁₈ O ₂ 170.25	14436-32-9	158 ²¹ ; 142 ⁴	4-02-00-01605 0.9238 ¹⁵	eth 4; EtOH 4 1.4507 ¹⁵
092	9-Decen-1-ol Decylenic alcohol	C ₁₀ H ₂₀ O 156,27	13019-22-2	236	4-01-00-02184 0.876 ²⁵	1.4480 ²⁰
093	3-Decen-2-one Heptylidene acetone	C ₁₀ H ₁₈ O	10519-33-2		4-01-00-03512	
094	1-Decen-3-yne	154.25 C ₁₀ H ₁₆ 136.24	33622-26-3	102-315.3	0.8473 ²⁰ 4-01-00-01105	1.4480 ²⁰
095	1-Decen-4-yne	136.24 C ₁₀ H ₁₈	24948-66-1	76 ²⁰	0.7873 ²⁰ 3-01-00-01049	1.4620 ²⁰
096	2-Decen-4-yne	136.24		73-4 ²²	0.7880 ²⁰	1.445 ²⁰
	•	C ₁₀ H ₁₆ 136.24	116668-40-7	55 ⁵	3-01-00-01049 0.7850 ²⁵	1.4609 ²⁵
097	1-Decyne Cctylacetylene	C ₁₀ H ₁₈ 138.25	764-93-2 -44	174	4-01-00-01054 0.7655 ²⁰	H ₂ O 1; EtOH 3; eth 3; os 3 1.4265 ²⁰
398	3-Decyne	C ₁₀ H ₁₈ 138.25	2384-85-2		4-01-00-01055	
999	4-Decyne	C ₁₀ H ₁₈ 138.25	2384-86-3	177	0.7619 ²⁵ 3-01-00-01017	1.4315 ²⁰
100	5-Decyne	138.25 C ₁₀ H ₁₈	1942-46-7	74.5 ¹⁹	0.772 ¹⁷ 4-01-00-01055	1.436 ¹⁷ H ₂ O 1; EtOH 3; eth 3
101	Dibutylacetylene 4-Decyne, 3,3-dimethyl-	138.25	-73	177; 78.8 ²⁵	0.7690 ²⁰	1.433120
102	3,3-Dimethyl-4-decyne	C ₁₂ H ₂₂ 168.31	70732-45-5	86 ²⁰	3-01-00-01026 0.7731 ²⁰	1,4399 ²⁰
	Deltamethrin Cyano(3-phenoxyphenyl)methyl-3-(2,2- dibromoethenyl)-2,2- dimethylcyclopropanecarboxylate	C ₂₂ H ₁₉ Br ₂ NO ₃ 505.21	52918-63-5 99	2869		
103	Demeton S methyl Phosphorothioic acid, S-[2-(ethylthio)ethyl] O,O-dimethyl ester	C ₈ H ₁₅ O ₃ PS ₂ 230.29	919-86-8	89 ^{0.15} ; 118 ¹	1.207 ²⁰	

PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS (continued)

No.	Name Synonym	Mol. Form. Mol. Wt.	CAS RN mp/°C	Merck No. bp/°C	Beil. Ref. den/g cm ⁻³	Solubility n _D
11619	Tetradecanenitrile Myristonitrile	C ₁₄ H ₂₇ N 209.38	629-63-0 19	226 ¹⁰⁰ ; 119 ¹	4-02-00-01139 0.8281 ¹⁹	H ₂ O 1; EtOH 5; eth 5; ace 5 1.4392 ²³
11620	1-Tetradecanesulfonic acid	C ₁₄ H ₃₀ O ₃ S 278.46	7314-37-6 65	110	4-04-00-00066 0.9996 ²⁵	H ₂ O 4
11621	Tetradecylsulfonic acid 1-Tetradecanethiol	C ₁₄ H ₃₀ S 230.46	2079-95-0	176-80 ²²	4-01-00-01867 0.8469 ²⁰	H ₂ O 1; EtOH 3; eth 3; ctc 3 1,4597 ²⁰
11622	Tetradecanoic acid	C ₁₄ H ₂₈ O ₂	544-63-8	6246 250 ¹⁰⁰	4-02-00-01126 0.8622 ⁵⁴	H ₂ O 1; EtOH 3; eth 2; ace 3 1.4723 ⁷⁰
11623	Myristic acid Tetradecanoic acid, anhydride	228.38 C ₂₈ H ₅₄ O ₃	53.9 626-29-9	250	4-02-00-01138	eth 4; EtOH 4 1,4335 ⁷⁰
11624	Tetradecanoic acid, ethyl ester	438.73 C ₁₆ H ₃₂ O ₂	53.4 124-06-1	205	0.8502 ⁷⁰ 4-02-00-01131 0.8573 ²⁵	H ₂ O 1; EtOH 3; eth 2; ctc 3 1,4362 ²⁰
11625	Tetradecanoic acid, methyl ester	256.43 C ₁₅ H ₃₀ O ₂ 242.40	12.3 124-10-7	295	4-02-00-01131	H ₂ O 1; EtOH 5; eth 5; ace 5 1.425 ⁴⁵
11626	Tetradecanoic acid, 1-methylethyl ester	C ₁₇ H ₃₄ O ₂	19 110-27-0	295; 155 ⁷ 5103	0.8671 ²⁰ 4-02-00-01132	H ₂ O 1; EtOH 3; eth 3; ace
11627	Isopropyl myristate Tetradecanoic acid, phenylmethyl ester	270.46 C ₂₁ H ₃₄ O ₂	31161-71-4	193 ²⁰ ; 140 ²	0.8532 ²⁰ 2-06-00-00417	1.4325 ²⁵ bz 4; eth 4; EtOH 4; chl 4
11628	Tetradecanoic acid, 1,2,3-propanetriyl ester	318.50 C ₄₅ H ₈₆ O ₆	20.5 555-45-3	229.3 ¹¹ 9638	0.9293 ²⁵ 4-02-00-01135	H ₂ O 1; EtOH 2; eth 3; ace 3
11629	Trimyństin Tetradecanoic acid, propyl ester	723.17 C ₁₇ H ₃₄ O ₂	56.5 14303-70-9	311	0.8848 ⁶⁰ 4-02-00-01132	1.4428 ⁶⁰ ace 4; bz 4; eth 4; EtOH 4
11630	1-Tetradecanol	270.46 C ₁₄ H ₃₀ O	112-72-1	147 ² 6248	0.8592 ²⁰ 4-01-00-01864	1.4356 ²⁵ H ₂ O 1; EtOH 4; eth 4; ace
11631	Tetradecyl alcoho! 2-Tetradecanol	214.39 C ₁₄ H ₃₀ O	39.5 4706-81-4	289	0.8236 ³⁸ 4-01-00-01867	
11632	3-Tetradecanol	214.39 C ₁₄ H ₃₀ O	34 1653-32-3	284	0.8315 ²⁰ 4-01-00-01868	1.4444 ²⁰ bz 4; eth 4; EtOH 4
11032	3-Tetradecanor	214.39	31.5	173 ²⁵ ; 146 ¹⁰	0.8098 ⁵³	1.4340 ⁴⁵
11633	2-Tetradecanone Dodecylmethylketone	C ₁₄ H ₂₈ O 212.38	2345-27-9 33.5	205 ¹⁰⁰ ; 134 ¹³	4-01-00-03389	H ₂ O 1; EtOH 3; ace 3; os 3
11634	3-Tetradecanone	C ₁₄ H ₂₈ O 212.38	629-23-2 34	152 ¹⁶	4-01-00-03389	H ₂ O 1; EtOH 3; ace 3; os 3
11635	Tetradecanoyl chloride Myristoyl chloride	C ₁₄ H ₂₇ CIO 246.82	112-64-1 -1	171 ¹⁶	4-02-00-01138 0.9078 ²⁵	eth 3
11636	1-Tetradecene	C ₁₄ H ₂₈ 196.38	1120-36-1 -12	233	4-01-00-00924 0.7745 ²⁵	H ₂ O 1; EtOH 4; eth 4; bz 3 1.4351 ²⁰
11637	4-Tetradecenoic acid	C ₁₄ H ₂₆ O ₂	544-65-0 18.5	185-8 ¹³	4-02-00-01626 0.9024 ²⁰	bz 4; peth 4 1,4559 ²⁰
11638	Tuduic acid 5-Tetradecenoic acid	226.36 C ₁₄ H ₂₆ O ₂	544-66-1	190-5 ¹⁵	3-02-00-01373 0.9046 ²⁰	1,4552 ²⁰
11639	Physoteric acid 9-Tetradecenoic acid	226.36 C ₁₄ H ₂₆ O ₂	20 13147-06-3	1440.6	4-02-00-01626	1,4519 ²⁰
11640	2-Tetradecyne	226,36 C ₁₄ H ₂₆ 194.36	-4 638-60-8		0.9018 ²⁰ 0-01-00-00262	eth 4; EtOH 4
11641	7-Tetradecyne	C ₁₄ H ₂₆	6.5 35216-11-6	252.5	0.8000 ²⁰ 3-01-00-01027	eth 4; EtOH 4
11642	2,5,8,11-Tetraoxadodecane	194.36 C ₈ H ₁₈ O ₄	112-49-2	144 ³⁰ 9604	0.7991 ²⁰ 4-01-00-02401	1.4330 ²⁵ H ₂ O 4; bz 4
11643	Triglyme 2,4,8,10-Tetraoxaspiro[5.5]undecane	178.23 C ₇ H ₁₂ O ₄	-45 126-54-5	216	0.986 ²⁰ 5-19-11-00342	1.4224 ²⁰ H ₂ O 4; ace 4; eth 4; EtOH
11644	Tetraphenylene	160.17 C ₂₄ H ₁₆	48.3 212-74-8	147 ⁵³ ; 68 ¹	4-05-00-02773	EtOH 3; eth 2; AcOEt 3;
,,,,,,	Tetrabenzocyclooctatetraene	304.39	233	sub 200		PhNO ₂ 3
11645	Tetraphosphoric acid, hexaethyl ester Ethyl tetraphosphate	C ₁₂ H ₃₀ O ₁₃ P ₄ 506.26	757-58-4 -40	150 dec	1.2917 ²⁷	ace 4; bz 4; EtOH 4 1,4273 ²⁷
11646	Tetrasiloxane, decamethyl- Decamethyltetrasiloxane	C ₁₀ H ₃₀ O ₃ Si ₄ 310.69	141-62-8 -76	2843 194	4-04-00-04119 0.8536 ²⁵	H ₂ O 1; EtOH 2; bz 3; peth 1,3895 ²⁰
11647	Tetrasiloxane, 1,1,1,3,5,7,7,7-octamethyl-	C ₈ H ₂₆ O ₃ Si ₄	16066-09-4	170	4-04-00-04098 0.8559 ²⁰	1.3854 ²⁰
11648	1,1,1,3,5,7,7,7-Octamethyltetrasiloxane Tetrasul	282.63 C ₁₂ H ₈ Cl ₄ S	2227-13-6	170	0.0009	1.0004
11649	ρ-Chlorophenyl 2,4,5-trichlorophenyl sulfide Tetrasulfide, bis(1,1-dimethylethyl)	324.06 C ₈ H ₁₈ S ₄	5943-35-1	700.2	4-01-00-01638	4.500020
11650	Di- <i>tert</i> -Butyl tetrasulfide Tetratriacontane	242,49 C ₃₄ H ₇₀	2.3 14167-59-0	70 ^{0.2}	1.0690 ²⁰ 4-01-00-00597	1.5660 ²⁰
11651	1,2,4,5-Tetrazine	478.93 C ₂ H ₂ N ₄	72.6 290-96-0	285.4 ³	0.7728 ⁹⁰ 4-26-00-01710	1.4296 ⁹⁰ H ₂ O 3; EtOH 3; eth 3; sulf
11652	sym-Tetrazine 1H-Tetrazole	82.06 CH ₂ N₄	99 288-94-8	sub	4-26-00-01652	H ₂ O 2
11653	2H-Tetrazolium, 2,3,5-triphenyl-, chloride	70.05 C ₁₉ H ₁₅ ClN ₄	157 298-96-4	sub 9658	1.4060 ²⁰ 4-26-00-01774	H ₂ O 3; EtOH 3; eth 1; ace
	Triphenyltetrazolium chloride 5H-Tetrazolo[1,5-a]azepine, 6,7,8,9-	334.81	243 dec 54-95-5	7097	4-26-00-01712	H ₂ O 4; EtOH 4; eth 3; ace
11654	tetrahydro-	C ₆ H ₁₀ N ₄ 138.17	59.5	194 ¹²	20-00-01712	1120 4, 21011 4, 001 -1
11655	Pentylenetetrazole 4-Thia-1-azabicyclo[3.2.0]heptane-2- carboxylic acid, 3,3-dimethyl-7-oxo-6-	C ₁₆ H ₁₇ N ₂ NaO ₄ S	69-57-8	1157	4-27-00-05861	
	[(phenylacetyl)amino]- Penicillin G, sodium salt	356.38				

PHYSICAL CONSTANTS OF ORGANIC COMPOUNDS (continued)

	Name Synonym	Mol. Form. Mol. Wt.	CAS RN mp/°C	Merck No. bp/°C	Beil, Ref. den/g cm ⁻³	Solubility n _D
	Dodecanoic acid, 2,3-dihydroxypropyl ester,	C ₁₅ H ₃₀ O ₄	40738-26-9		4-02-00-01096	EtOH 2; eth 4; ace 4; bz 3
	(±)- Glycerol, 1-laurate (<i>DL</i>)	274.40	63	186 ²	0.9248 ⁹⁷	1.4350 ⁸⁶ eth 4; EtOH 4
5311	Dodecanoic acid, 1,2-ethanediyl ester	C ₂₆ H ₅₀ O ₄ 426.68	624-04-4	4393 188 ²⁰	4-02-00-01094	
	Ethylene glycol dilaurate		56.6 106-33-2	3774	4-02-00-01092	H ₂ O 1; EtOH 4, eth 5; ctc 2
5312	Dodecanoic acid, ethyl ester Ethyl laurate	C ₁₄ H ₂₈ O ₂ 228.38	-10	271; 154 ¹⁵	0.8618 ²⁰	1.4311 ²⁰ EtOH 5; eth 5; ace 5; bz 3
5313	Dodecanoic acid, 2-(2-hydroxyethoxy)ethyl	C ₁₆ H ₃₂ O ₄	141-20-8	3110		EtOH 5, eth 5, ace 5, bz 5
0010	ester	288.43	17.5	>270	0.96 ²⁵	
504.4	Diethylene glycol monolaurate Dodecanoic acid, 2-methyl-	C ₁₃ H ₂₆ O ₂	2874-74-0	1	4-02-00-01121 0.890 ¹⁸	
5314	Dodecanoic acid, 2-metry:	214.35	22 111-82-0	1531	4-02-00-01090	H ₂ O 1; EtOH 5; eth 5; ace 5
5315	Dodecanoic acid, methyl ester	C ₁₃ H ₂₆ O ₂ 214.35	111-82-0 5.2	267	0.8702 ²⁰	1.4319 ²⁰
	Methyl laurate Dodecanoic acid, 1-methylethyl ester	C ₁₅ H ₃₀ O ₂	10233-13-3	607	4-02-00-01092 0.8536 ²⁰	eth 4; EtOH 4 1,4280 ²⁵
5316	Dodecarioic acid, i-metrylettyl ester	242.40		196 ⁶⁰ ; 117 ²	4-06-00-00618	ace 4; eth 4; EtOH 4
5317	Dodecanoic acid, phenyl ester	C ₁₈ H ₂₈ O ₂ 276.42	4228-00-6 24.5	210 ¹⁵	0.935430	
	Phenyl laurate	C ₁₉ H ₃₀ O ₂	140-25-0		4-06-00-02267	bz 4; eth 4; EtOH 4; peth 4
5318	Dodecanoic acid, phenylmethyl ester	290.45	8.5	209-11 ¹²	0.9429 ²⁵ 4-02-00-01098	1.4812 ²⁴ H ₂ O 1; EtOH 3; eth 3; ace
5319	Dodecanoic acid, 1,2,3-propanetriyl ester	C ₃₉ H ₇₄ O ₆	538-24-9		0.898655	1.4404 ⁶⁰
		639.01 CasHaoOa	3681-78-5		4-02-00-01092	1.4335 ²⁰
5320	Dodecanoic acid, propyl ester Propyl laurate	C ₁₅ H ₃₀ O ₂ 242.40		205 ⁸⁰ ; 124 ²	0.8600 ²⁰ 4-01-00-01844	H ₂ O 1; EtOH 3; eth 3; bz 2
5321	1-Dodecanol	C ₁₂ H ₂₆ O	112-53-8 24	3402 259	0.830924	1120 1, 21011 0, 0111 1, 1
	Lauryl alcohol	186.34 C ₁₂ H ₂₈ O	10203-28-8	200	3-01-00-01793	
5322	2-Dodecanol	186.34	19	252	0.8286 ²⁰	1.4400 ²⁰
5323	3-Dodecanol	C12H26O	10203-30-2	130 ¹⁵	4-01-00-01854 0.8223 ³²	
3323	0-D00000	186.34	25 6836-38-0	130**	3-01-00-01794	eth 4; EtOH 4
5324	6-Dodecanol	C ₁₂ H ₂₆ O 186.34	30	225; 119 ⁹	0.820140	H ₂ O 1; EtOH 3; eth 3; ace
E225	2-Dodecanone	C ₁₂ H ₂₄ O	6175-49-1	046.5	4-01-00-03382 0.8198 ²⁰	1,4330 ²⁰
5325	Decyl methyl ketone	184.32	21 6064-27-3	246.5	4-01-00-03383	
5326	6-Dodecanone	C ₁₂ H ₂₄ O 184.32	10	1129		1.4302 ²⁰
5007	Amyl hexyl ketone 1-Dodecanone, 1-phenyl-	C18H28O	1674-38-0	05	4-07-00-00847 0.8794 ¹⁸	H ₂ O 1; ace 3; ctc 2 1.4700 ¹⁸
5327	1-Dodecanone, 1-phenyl	260.42	47	201 ⁹ , 181 ⁵	4-02-00-01103	eth 4
5328	Dodecanoyl chloride	C ₁₂ H ₂₃ CIO 218.77	112-16-3 -17	145 ¹⁸	0.9169 ²⁵	1.4458 ²⁰
	1,3,6,10-Dodecatetraene, 3,7,11-trimethyl-,	C ₁₅ H ₂₄	502-61-4	3883	3-01-00-01067	H ₂ O 1; eth 3; ace 3; peth
5329	(<i>E.E</i>)-			129-3212	0.8410 ²⁰	1.4836 ²⁰
	α-Famesene	204.36	19317-11-4	129-52	4-01-00-03603	
5330	2,6,10-Dodecatrienal, 3,7,11-trimethyl-	C ₁₅ H ₂₄ O 220.35	19517-11-4	172-4 ¹⁴	0.893 ¹⁸	1.4995 ace 4; eth 4; chl 4
5331	1,6,10-Dodecatriene, 7,11-dimethyl-3-	C ₁₅ H ₂₄	18794-84-8	3884	4-01-00-01133	ace 4; eth 4, chi 4
5331	methylene-, (E)-			121-2 ⁹	0.8363 ²⁰	1.4899 ²⁰
	β-Farnesene	204.38 C ₁₅ H ₂₆ O	142-50-7	6388	4-01-00-02336	EtOH 4; eth 3; ace 3; os
5332	1,6,10-Dodecatrien-3-ol, 3,7,11-trimethyl-, [S-	01511260		0.1	0.8778 ²⁰	1,4898 ²⁰
	(Z)]-	222.37	100.00.5	276; 70 ^{0.1}	4-01-00-02335	H ₂ O 1; EtOH 4; eth 3; ac
5333	2,6,10-Dodecatrien-1-ol, 3,7,11-trimethyl-,	C ₁₅ H ₂₆ O	106-28-5			•
	(É,E)-	222.37		160 ¹⁰	0.8846 ²⁰	1.4877 ²⁰ ace 4; eth 4; EtOH 4
F224	2,6,10-Dodecatrien-1-ol, 3,7,11-trimethyl-,	C ₁₅ H ₂₆ O	3790-71-4		4-01-00-02335	ace 4, eth 4, Eton 4
5334	(Z,E)-	222.37		156 ¹² ;	0.8908 ²⁰	1.4877 ²⁰
		222.31		1200.3		H ₂ O 1; EtOH 3; eth 3; ac
5335	1-Dodecene	C ₁₂ H ₂₄	112-41-4	213.8	4-01-00-00914 0.7584 ²⁰	1.4300 ²⁰
5335		168.32	-35.2	9493	4-02-00-02279	eth 4; EtOH 4; chl 4
5336	2-Dodecenedioic acid, (E)-	C ₁₂ H ₂₀ O ₄ 228.29	6402-36-4 165.5	5-55		
	Traumatic acid 2-Dodecenoic acid	C12H22O2	4412-16-2	13	4-02-00-01619 0.9265 ²⁰	1.4629 ²⁵
5337	Z-Dodecenoic acid	C ₁₂ H ₂₂ O ₂ 198.31	17.1	155 ³ ; 127 ^{0,15}	U.920022	
		C . U . O	505-92-0		4-02-00-01619	bz 4; eth 4; chl 4
5338	4-Dodecenoic acid	C ₁₂ H ₂₂ O ₂ 198.31	1.3	171 ¹³	0.9081 ¹⁵	1.4529 ²⁰
5339	Lindenc acid 5-Dodecenoic acid	C ₁₂ H ₂₂ O ₂	2761-84-4	170-2 ¹³	4-02-00-01619 0.9081 ²⁰	
5338		198.31	1.3 65423-25-8	170-2	4-02-00-01618	20
5340	11-Dodecenoic acid	C ₁₂ H ₂₂ O ₂ 198.31	20	171 ¹³ ; 144 ³		1.4510 ²⁰
- 5044	11-Dodecenoic acid, methyl ester	C ₁₃ H ₂₄ O ₂	29972-79-0	400.013	4-02-00-01618 0.8789 ²²	1.4414 ²⁰
5341	Methyl 11-dodecenoate	212.33	74744 26 6	136-9 ¹³	4-01-00-01112	
5342	1-Dodecen-3-yne	C ₁₂ H ₂₀ 164.29	74744-36-8	78 ⁴	0.7858 ²⁵	1.4510 ²⁵
	4 Dadama	C ₁₂ H ₂₂	765-03-7		4-01-00-01066 0.7788 ²⁰	1.4340 ²⁰
5343	1-Dodecyne Decylacetylene	166.31	-19	215	0-01-00-00261	
5344	2-Dodecyne	C ₁₂ H ₂₂ 168.31	629-49-2 -9	105 ¹⁵	0.7917 ¹⁵	1.482820
		C ₁₂ H ₂₂ 166.31	6790-27-8		3-01-00-01025	ace 4; eth 4
5345	3-Dodecyne	~12' <u>'22</u>		9512	0.7871 ²⁰	1.4442 ²⁰